(21) International Application Number:

## WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau





### INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 5:		(11) International Publication Numbe	r: WO 92/15673
C12N 9/02, 15/00, 15/11 C12N 15/53, 15/70, 15/74	A1	(43) International Publication Date:	17 September 1992 (17.09.92)
			-

PCT/US91/01614

(22) International Filing Date: 11 March 1991 (11.03.91)

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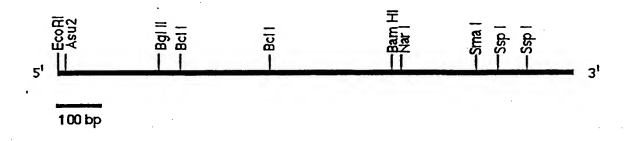
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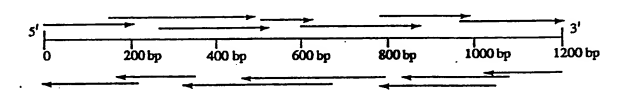
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#### Published

With international search report.

#### (54) Tide: CLONING AND EXPRESSION OF RENILLA LUCIFERASE





#### (57) Abstract

Genetic material encoding luciferase from the marine coelenterate Renilla has been isolated and characterized. This genetic material allows the production of peptides for use as labels in bioluminescence assays or can itself be directly used to identify luciferase genes from related organisms.

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## CLONING AND EXPRESSION OF RENILLA LUCIFERASE

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### BACKGROUND OF THE INVENTION

#### Field of the Invention

This invention relates to the field of genetic 10 engineering and is particularly related to the expression of proteins by techniques involving genetic engineering.

#### 15 Description of the Background

The Renilla, also known as sea pansies, belong to a class of coelenterates known as the anthozoans. In addition to Renilla, other representative bioluminescent genera of the class Anthozoa include Cavarnularia, Ptilosarcus, Stylatula, Acanthoptilum, and Parazoanthus. All of these organisms are bioluminescent and emit light as a result of the action of an enzyme (luciferase) on a substrate (luciferin) under appropriate biological conditions. Prior studies have demonstrated that all of the above-mentioned anthozoans contain similar luciferases and luciferins. See, for example, Cormier et al., J. Cell. Physiol. (1973) 81: 291-298. The luciferases and luciferins from each of these anthozoans will cross-

react with one another to produce the characteristic 30 blue luminescence observed in Renilla extracts. Each of these luciferases has similar biochemical properties, and the biochemical requirements for bioluminescence are identical regardless of the anthozoan from which the luciferase was derived. 35

There has been considerable interest of late in replacing radioactive labels used in analytical

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assays with other types, such as luminescent labels. Firefly luciferase, which is a molecule of significantly different structure that does not react with Renilla-like luciferins, is one molecule that has been proposed for use as such labels. However, firefly luciferase suffers from a number of deficiencies that make this molecule less than optimal in biological assays. For example, ATP acts as a trigger of the firefly luciferase system, and the ubiquitous nature of ATP makes control of this variable difficult.

A prior patent application by one of the present inventors, U.S. Patent Application Serial No. 059,137, filed June 5, 1987, describes use of coelenterate-derived luciferases and photoproteins as bioluminescent labels. Other applications by the same inventor, for example, U.S. Application Serial Nos. 173,045, filed March 17, 1988, and 165,422, filed February 29, 1988, describe recombinant DNA capable of expressing the photoprotein apoaequorin.

20 The photoprotein aequorin (which consists of apoaequorin bound to a coelenterate luciferin molecule) and Renilla luciferase both utilize the same coelenterate luciferin, and the chemistry of light emission in both cases has been shown to be the same. However, aequorin luminescence is triggered by calcium, 25 does not require dissolved oxygen, and represents a single turnover event. In contrast, Renilla luciferase is not triggered by calcium and requires dissolved oxygen in order to produce light in the presence of 30 coelenterate luciferin. Renilla luciferase also acts as a true enzyme, catalyzing a long-lasting luminescence in the presence of saturating levels of luciferin.

Sub-attomole levels of aequorin can be

detected with photometers even though its luminescence represents a single turnover event. Renilla luciferase, because of its enzymatic ability, should be

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detectable at levels 1 to 2 orders of magnitude lower than aequorin. Furthermore, Renilla luciferase is known to be relatively stable to heat, an important consideration for assays that often involve incubation at physiological temperatures. Accordingly, Renilla luciferase is a potentially useful label for biological and other assays.

On the other hand, Renilla live on the ocean bottom, about 30 to 100 feet deep, and must be collected by dregging. From 1 kg of Renilla (about 1000 animals), approximately 1 mg of pure Renilla luciferase can be obtained following a tedious procedure which requires purifying the protein about 12,000 fold. The purification procedure is described in Matthews et al., Biochemistry (1977) 16: 85-91. As a result, there has been no development of Renilla luciferase as a detectable label.

Accordingly, improved techniques for the production of pure Renilla luciferase are necessary before this molecule can be used commercially in bioluminescence assays.

## BRIEF DESCRIPTION OF THE DRAWINGS

The invention will be better understood by
reference to the following detailed description and
examples and the attached Figures that form part of the
present specification, wherein:

Figure 1 is the nucleotide sequence of a clone that contains a Renilla reniformis luciferase cDNA sequence.

Figure 2 is the amino acid sequence derived from the open reading frame of the Renilla luciferase cDNA shown in Figure 1.

Figure 3 is the recombinant luciferase amino acid sequence with different types of underlining to show the location of peptides obtained by digestion of native luciferase with V-8 protease.

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Figure 4 is a table showing the amino acid sequence of Renilla reniformis peptides obtained by V-8 protease digestion. Regions of low degeneracy that were selected for preparation of oligonucleotide probes are shown by boxes. The probes are shown at the bottom part of the Figure.

Figure 5 is a schematic representation of a restriction enzyme map for Renilla luciferase cDNA. The lower portion of Figure 5 is a schematic representation of sequencing strategy for Renilla luciferase cDNA.

Figure 6 is a map of a Renilla luciferase expression plasmid.

Figure 7 is a schematic diagram of the control 15 region of the plasmid pTZRLuc-l.

Figure 8 is a schematic diagram of the purification scheme used to purify recombinant luciferase.

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## SUMMARY OF THE INVENTION

The present invention provides genetic material encoding Renilla luciferase. The genetic material can be used to produce the enzyme for use as luminescent tags in bioluminescence assays and for 25 other purposes for which such labels are desirable. Additionally, the genetic material can be used as a source of probes that can be used in nucleic acid hybridization assays for the identification of other luciferase genes from related organisms. Fragments of the enzyme can be used to prepare antibodies for the purpose of identifying luciferase genes from related organisms. Specific genetic materials and luciferase proteins are disclosed in the following detailed description and examples.

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## DESCRIPTION OF SPECIFIC EMBODIMENTS

The present inventors have identified and obtained for the first time genetic material encoding luciferase from the coelenterate genus Renilla previously has been available only in limited 5 quantities. Since luciferases have a number of uses as a bioluminescent label and since Renilla luciferase has a number of properties that make it particularly useful as a label, availability of the enzyme in significant quantities in pure form provides a significant 10 commercial advantage over prior sources. The Renilla genetic material also provides a source of nucleic acid probes for use in hybridization techniques that allow location of luciferase genes in related organisms. cDNA sequence for a clone that contains a Renilla 15 reniformis luciferase gene is set forth in Figure 1, with the translated cDNA amino acid sequence being set forth in Figure 2. The coding sequence of the clone in Figure 1 begins at nucleotide 10 and continues to a 20

stop codon at nucleotide 944. Figure 3 shows a complete recombinant Renilla luciferase amino acid sequence as produced by an expression system.

The present invention has specifically contemplated each and every possible variation of polynucleotide that could be made by selecting 25 combinations based on the possible codon choices listed in Figure 1 (with the reading frame beginning at position 1 of Figure 1) and in Table 1 (below), and all such variations are to be considered as being

specifically disclosed and equivalent to the sequence of Figure 1. Codons are preferably selected to fit the host cell in which the enzyme is being produced, Selection of codons to maximize expression of proteins in a heterologous host is a known technique.

35 Other DNA molecules that code for such peptides can readily be determined from the list of codons in Table 1 and are likewise contemplated as being equi-

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valent to the DNA sequence of Figure 1. In fact, since there is a fixed relationship between DNA codons and amino acids in a peptide, any discussion in this application of a replacement or other change in a peptide is equally applicable to the corresponding DNA sequence or to the DNA molecule, recombinant vector, or transformed microorganism in which the sequence is located (and vice versa).

### TABLE 1

	GENETIC CODE
Alanine (Ala, A)	GCA, GCC, GCG, GCT
Arginine (Arg, R)	AGA, AGG, CGA, CGC, CGG, CG
Asparagine (Asn, N)	AAC, AAT
Aspartic acid (Asp, D)	GAC, GAT
Cysteine (Cys, C)	TGC, TGT
Glutamine (Gln, Q)	CAA, CAG
Glutamic acid (Glu, E)	GAA, GAG
Glycine (Gly, G)	GGA, GGC, GGG, GGT
Histidine (His, H)	CAC, CAT
Isoleucine (Ile, I)	ATA, ATC, ATT
Leucine (Leu, L)	
Lysine (Lys, K)	CTA, CTC, CTG, CTT, TTA, TTG AAA, AAG
Methionine (Met, M)	ATG
Phenylalanine (Phe, F)	TTC, TTT
Proline (Pro, P)	CCA, CCC, CCG, CCT
Serine (Ser, S)	
Threonine (Thr, T)	AGC, AGT, TCA, TCC, TCG, TCT ACA, ACC, ACG, ACT
Tryptophan (Trp, W)	TGG
Tyrosine (Tyr, Y)	TAC, TAT
Valine (Val, V)	
Termination signal	GTA, GTC, GTG, GTT
	TAA, TAG, TGA

Key: Each 3-letter triplet represents a trinucleotide of DNA having a 5' end on the left and a 3' end on the right. The letters stand for the purine or pyrimidine bases forming the nucleotide sequence: A = adenine, G = guanine, C = cytosine, and T = thymine. The RNA code is the same except that U (uracil) replaces T.

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In addition to the specific nucleotides listed in Figure 1, DNA (or corresponding RNA) molecules of the invention can have additional nucleotides preceding or following those that are specifically listed. For example, poly A can be added to the 3'-terminal; a 5 short (e.g., fewer than 20 nucleotides) sequence can be added to either terminal to provide a terminal sequence corresponding to a restriction endonuclease site, stop codons can follow the peptide sequence to terminate 10 translation, and the like. Additionally, DNA molecules containing a promoter region or other control region upstream from the gene can be produced. All DNA molecules containing the sequences of the invention will be useful for at least one purpose since all can minimally be fragmented to produce oligonucleotide probes and be 15 used in the isolation or detection of DNA from biological sources.

A number of words used in this specification have specific meanings in addition to their more common meanings. "Renilla luciferase" means the luciferase 20 enzyme isolated from a member of the genus Renilla or an equivalent molecule obtained from any other source or synthetically. By "equivalent" is meant, when referring to two nucleotide sequences, that the two nucleotide sequences in question encode the same 25 sequence of amino acids. When "equivalent" is used in referring to two peptides, it means that the two peptides will have substantially the same amino acid sequence. When "equivalent" refers to a property, the property does not need to be present to the same extent (e.g., two peptides can exhibit different rates of the same type of enzymatic activity), but the properties are preferably substantially the same. "Complementary," when referring to two nucleotide sequences, means that the two sequences are capable of 35 hybridizing, preferably with less than 25%, more preferably with less than 15%, even more preferably

with less than 5%, most preferably with no mismatches between opposed nucleotides. Preferred hybridizing conditions (which are not limited to specific numbers of mismatches) are set forth in the Examples. "substantially" varies with the context as understood by those skilled in the relevant art and generally means at least 70%, preferably means at least 80%, more preferably at least 90%, and most preferably at least The phrase "substantially identical" includes complete identity as well as less than complete 10 identity (e.g., of amino acid sequences or enzymatic activity) as established by the prior definition of "substantially." The term "isolated" as used herein refers to, e.g., a peptide, DNA, or RNA separated from other peptides, DNAs, or RNAs, respectively, and being 15 found in the presence of (if anything) only a solvent, buffer, ion or other component normally present in a biochemical solution of the same. "Isolated" does not encompass either natural materials in their native state or natural materials that have been separated 20 into components (e.g., in an acrylamide gel) but not obtained either as pure substances or as solutions. The phrase "replaced by" or "replacement" as used herein does not necessarily refer to any action that must take place but to the peptide that exists when an 25 indicated "replacement" amino acid is present in the same position as the amino acid indicated to be present in a different formula (e.g., when leucine instead of valine is present at amino acid 11).

Since the DNA sequence of the Renilla luciferase gene has been identified, it is possible to produce a DNA gene entirely by synthetic chemistry, after which the gene can be inserted into any of the many available DNA vectors using known techniques of recombinant DNA technology. Thus, the present invention can be carried out using reagents, plasmids, and microorganisms which are freely available and in the

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public domain at the time of filing of this patent application without requiring a deposit of genetic material.

- For example, nucleotide sequences greater than 100 bases long can be readily synthesized on an Applied Biosystems Model 380A DNA Synthesizer as evidenced by commercial advertising of the same (e.g., Genetic Engineering News, November/December 1984, p. 3). Such oligonucleotides can readily be spliced using, among
- others, the technique of preparing overlapping complementary sequences (e.g, 1-100 of coding strand, 0-50 and 51-150 of complementary strand, 101-200 of coding strand, etc.), followed by hybridizing and ligating the strands. Such techniques are well known and are
- described in detail in, for example, Davis et al.,

  Basic Methods in Molecular Biology, Elsevier Science
  Publ. Co., Inc., New York (1986). The peptides can
  then be expressed in a host organism as described
  herein.
- Furthermore, automated equipment is also available that makes direct synthesis of many of the peptides disclosed herein readily available, especially peptide fragments of less than the entire Renilla luciferase enzyme. In the same issue of Genetic

  Engineering News mentioned above a particular in the same issue of Genetic
- 25 Engineering News mentioned above, a commercially available automated peptide synthesizer having a coupling efficiency exceeding 99% is advertised (page 34). Such equipment provides ready access to the peptides of the invention, either by direct synthesis or by synthesis of a series of fragments that can be coupled using other known techniques.

In addition to the specific polypeptide sequence shown in Figures 2 and 3, peptide fragments based on these sequences and fragments and full length sequences representing minor variations thereof will have at least some of the biological activities of luciferase and will therefore be useful in appropriate

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circumstances. For example, fragments of the luciferase enzyme sequence can readily be prepared and can be screened for use as luciferin binding site models. Peptide synthesizers can be used to prepare small polypeptide fragments (e.g., less than 100 amino acids) or techniques of genetic engineering can be used to prepare larger fragments. A simple screening procedure that will identify suitable polypeptide fragments consists of attaching a suitable substrate, e.g., a coelenterate luciferin molecule, to an affinity column and capturing peptide fragments that are retained by the bound substrate. Such peptides can also be used (and are indeed more likely to be used) as immunogens for the preparation of antibodies that can be used to screen for the expression of a luciferase by

a genetically engineered organism, in which case the bound substrate will be an antibody or similar molecule that binds specifically to Renilla luciferase.

The ability to prepare and select peptide
fragments having appropriate binding affinity from a larger protein is well trans-

fragments having appropriate binding affinity from a larger protein is well known in the art and is described in a number of publications, including patents. See, for example, U.S. Patent No. 4,629,783, which describes the preparation of immunologically active fragments of viral proteins that bind with the same antibodies as the entire viral protein.

In addition, minor variations of the previously mentioned peptides and DNA molecules are also contemplated as being equivalent to those peptides and DNA
molecules that are set forth in more detail, as will be
appreciated by those skilled in the art. For example,
it is reasonable to expect that an isolated replacement
of a leucine with an isoleucine or valine, an aspartate
with a glutamate, a threonine with a serine, or a similar replacement of an amino acid with a structurally
related amino acid (i.e., a conservative replacement)
will not have a major effect on the biological activity

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of the resulting molecule, especially if the replacement does not involve an amino acid at a binding site or other site of biologic activity. Furthermore, additional amino acids can be present at either of the two termini, or amino acids can be absent from one or both of the termini, as is known in the art.

Whether a change results in a functioning peptide can readily be determined by direct analysis for function in a assay that relies on ability of the modified enzyme (or fragment) to carry out the normal 10 function of the natural luciferase enzyme (or fragment). For example, modified peptides can be tested for ability to catalyze the emission of light from coelenterate luciferin by the same techniques described below for the recombinant Renilla luciferase 15 molecule. Peptides in which more than one replacement has taken place can readily be tested in the same manner. Preferred peptides differ at no more than 12, more preferably no more than 5, amino acids in any contiguous group of 20 amino acids. Substitutions of 20 amino acids, when they occur, are preferably from within standard conservative groups. Standard conservative groups of amino acids are shown in parenthesis using the one-letter amino acid code: polar (A,V,L,I,P,M); aromatic (F,T,W); uncharged polar 25 (G,S,T,C,N,Q); acidic (D,E); basic (K,R,H). The aromatic amino acids are sometimes considered to belong to the broader-defined nonpolar (F,W) or uncharged polar (T) groups.

Salts of any of the peptides described herein will naturally occur when such peptides are present in (or isolated from) aqueous solutions of various pHs. All salts of peptides having the indicated biological activity are considered to be within the scope of the present invention. Examples include alkali, alkaline earth, and other metal salts of carboxylic acid residues, acid addition salts (e.g., HCl) of amino residues.

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dues, and zwitterions formed by reactions between carboxylic acid and amino residues within the same molecule.

Although genes and corresponding proteins can be prepared by the totally synthetic techniques dis-5 cussed above, in preferred embodiments of the invention genetic information is obtained from natural sources and identified as described herein. The genetic material is first obtained in the form of a gene library, using any of numerous existing techniques. 10 of these is to randomly shear genomic DNA and insert this sheared material into expression vectors. enough recombinants are generated, there is a good probability of having at least one recombinant in the population which is expressing a fusion protein corre-15 sponding to the enzyme of interest.

Another strategy for preparing gene libraries is to make complementary DNA (cDNA) copies of the total mRNA population of the organism and to clone these as recombinant molecules in expression vectors. The expected nature of the organism (i.e., it was expected to have the characteristics of a eucaryote) indicated that introns might be present within the coding region of the desired gene. Although introns do not preclude use of sheared genomic DNA, they increase the number of recombinants which must be screened and make further analyses substantially complicated. Based on this result, use of a cDNA library to obtain Renilla genes is preferred.

Such a library was generated in the laboratory of the inventors and screened for expression of a gene product having luciferase activity. Details of this example are set forth below, including details of the experiments that lead to obtaining the complete sequence of the gene. However, there is no reason to believe that the sequence and specific engineered organism prepared by the inventors is any better than

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other clones that can be prepared using the guidance set forth in this specification. In fact, it is likely that expression of Renilla luciferase can be enhanced over that described herein by selection of other expression systems, as discussed in the examples below.

Now that the sequence of Renilla luciferase has been determined, it is no longer necessary to go through these steps to obtain the genetic material of the present invention. The polymerase chain reaction (PCR) technique can now be used to isolate genes from natural sources in a simpler and more direct manner. The PCR technique, including use in diagnosis, is disclosed in U.S. Patent 4,683,202, which is herein incorporated by reference. Since Renilla specimens are readily available from the oceans of the world, and since PCR probes can be prepared using the sequences set forth in this specification, it is possible to obtain any desired segment of the sequences set forth herein using the PCR technique and naturally available sources of Renilla genomic material. A specific example of such a technique for isolating the Renilla luciferase chromosomal gene is described in the examples that follow. The cloned gene can then be inserted into commercial vectors and expressed.

Although the techniques set forth above, when used in combination with the knowledge of those skilled in the art of genetic engineering and the previously stated guidelines, will readily enable isolation of the desired gene and its use in recombinant DNA vectors now that sufficient information is provided to locate the gene, other methods which lead to the same result are also known and may be used in the preparation of recombinant DNA vectors of this invention.

Expression of Renilla protein can be enhanced

by including multiple copies of the gene in a transformed host; by selecting a vector known to reproduce
in the host, thereby producing large quantities of pro-

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tein from exogeneous inserted DNA (such as pUC8; ptac12; pIN-III-ompA1, 2, or 3; pOTS; pAS1; or pKK223-3); or by any other known means of enhancing peptide expression.

5 One common variation is the preparation of a polypeptide of the invention in the form of a fused polypeptide. Such peptides are typically prepared by using the promoter region of a gene known to be expressed in a host and inserting nucleotides that encode all or a major portion of the amino acid sequence of 10 the invention into the genetic sequence for the host protein. Examples of such fused proteins include  $\beta$ galactosidase fused proteins. If desired, the fused peptide can be designed so that a site recognized by a proteolytic enzyme is present at the junction between 15 the two fused proteins. The proteolytic enzyme can then be used to cleave the expressed protein so that the desired luciferase enzyme is available in pure form.

In all cases, a <u>Renilla</u> luciferase will be expressed when the DNA sequence is functionally inserted into the vector. By "functionally inserted" is meant in proper reading frame and orientation, as is well understood by those skilled in the art. Typically, a gene will be inserted downstream from a promoter and will be followed by a stop codon, although production as a hybrid protein (possibly followed by cleavage) may be used, if desired.

In addition to the above general procedures
which can be used for preparing recombinant DNA molecules and transformed unicellular organisms in accordance with the practices of this invention, other known
techniques and modifications thereof can be used in
carrying out the practice of the invention. In particular, techniques relating to genetic engineering have
recently undergone explosive growth and development.
Many recent U.S. patents disclose plasmids, genetically

engineering microorganisms, and methods of conducting genetic engineering which can be used in the practice of the present invention. For example, U.S. Patent 4,273,875 discloses a plasmid and a process of isolating the same. U.S. Patent 4,304,863 discloses a pro-5 cess for producing bacteria by genetic engineering in which a hybrid plasmid is constructed and used totransform a bacterial host. U.S. Patent 4,419,450 discloses a plasmid useful as a cloning vehicle in recombinant DNA work. U.S. Patent 4,362,867 discloses 10 recombinant cDNA construction methods and hybrid nucleotides produced thereby which are useful in cloning processes. U.S. Patent 4,403,036 discloses genetic reagents for generating plasmids containing multiple copies of DNA segments. U.S. Patent 4,363,877 disclo-15 ses recombinant DNA transfer vectors. U.S. Patent 4,356,270 discloses a recombinant DNA cloning vehicle and is a particularly useful disclosure for those with limited experience in the area of genetic engineering since it defines many of the terms used in genetic 20 engineering and the basic processes used therein. U.S. Patent 4,336,336 discloses a fused gene and a method of making the same. U.S. Patent 4,349,629 discloses plasmid vectors and the production and use thereof. Patent 4,332,901 discloses a cloning vector useful in 25 recombinant DNA. Although some of these patents are directed to the production of a particular gene product that is not within the scope of the present invention, the procedures described therein can easily be modified to the practice of the invention described in this 30 specification by those skilled in the art of genetic engineering.

The implications of the present invention are significant in that useful amounts of Renilla

1 luciferase and genetic material of the invention will become available for use in the development of hybridization assays or in any other type of assay utilizing

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these materials. Transferring the Renilla luciferase cDNA which has been isolated to other expression vectors will produce constructs which improve the expression of luciferase in E. coli or express the polypeptide in other hosts.

Particularly contemplated is the isolation of genes from related organisms using oligonucleotide probes based on the principal and variant nucleotide sequences disclosed herein. Such probes can be considerably shorter than the entire sequence but should be at least 10, preferably at least 14, nucleotides in length. Intermediate oligonucleotides from 20 to 500, especially 30 to 200, nucleotides in length provide particularly specific and rapid-acting probes. Longer oligonucleotides are also useful, up to the full length of the gene. Both RNA and DNA probes can be used.

In use, the probes are typically labelled in a detectable manner (e.g., with 32p, 3H, biotin, or avidin) and are incubated with single-stranded DNA or RNA 20 from the organism in which a gene is being sought. Hybridization is detected by means of the label after single-stranded and double-stranded (hybridized) DNA (or DNA/RNA) have been separated (typically using nitrocellulose paper). Hybridization techniques suitable for use with oligonucleotides are well known.

Although probes are normally used with a detectable label that allows easy identification, unlabeled oligonucleotides are also useful, both as precursors of labeled probes and for use in methods that provide for direct detection of double-stranded DNA (or DNA/RNA). Accordingly, the term "oligonucleotide probe" refers to both labeled and unlabeled forms.

35 In summary, the inventors have reduced the present invention to practice by isolating and sequencing a cDNA clone for Renilla reniformis

luciferase. The deduced amino acid sequence from this cDNA, beginning at the first methionine residue, predicts a protein of M<sub>r</sub> equal to 36 kd, which is the approximate size of native Renilla luciferase. The deduced amino acid sequence also contains within it all six peptide sequences from V-8 protease-digested native Renilla luciferase. Only one mis-match was foundbetween these two sets of amino acid data, a substitution of a tryptophan for a leucine present in the peptide sequence. Comparisions of the native amino acid composition and the predicted recombinant luciferase composition reveal a very high degree of similarity with many identities between specific amino acid residues.

15 Additionally, expression of luciferase in a genetically engineered organism has been demonstrated. Luciferase activity was found in crude extracts of the original luciferase clone \alpha RLuc-6. Subcloning the cDNA into the vector pTZ18R increased this activity enough to allow the purification of 20 recombinant luciferase from the pTZRLuc-1 cells. Recombinant luciferase can be purified by a much simplified method from that previously used in the purification of native luciferase. The recombinant luciferase functions identically to native luciferase 25 in all aspects analysed thus far. Like native, recombinant luciferase has an emission spectrum with a  $\lambda$ max at 480 nm and a shoulder at 400 nm. absorption spectrum of recombinant luciferase is also identical to that of native. Additionally, both native 30 and recombinant luciferase are very stable at 37°C for several hours as well as having significant stability at 45°C. Using the specific activity determined for native luciferase, protein determinations made based on light emission correlate very well with  ${\rm A}_{280}$  and Lowry 35 protein determinations, suggesting that the specific activity of recombinant luciferase is similar to, if

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not the same as, that of native luciferase. Finally, amino-terminus amino acid sequence analysis of recombinant luciferase shows an identical sequence to that of the cDNA-predicted amino acid sequence from residues 2 through 18. A significant amount of the recombinant protein is blocked at the amino terminus, probably by N-formyl methionine, which accounts for the inability to determine the amino acid at residue 1.

The invention now being generally described, the same will be better understood by reference to the following examples which are provided for purposes of illustration only and are not to be considered limiting of the invention unless so specified.

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#### **EXAMPLES**

## Assay for Luciferase Activity

Crude supernatants or pure recombinant luciferase samples {10 to 100 µl} were added to 1 ml of luciferase assay buffer {0.5M NaCl, 0.1M KPO4 pH 7.6, 1mM EDTA, 0.02% BSA, and 0.004% NaN3} and vortexed in 12 X 75 mm test tube. Synthetic benzyl luciferin {10µl of a 2.5 nmol/µl stock} was added to the reaction to give a final concentration of 2.5 X 10<sup>-8</sup> M, and the mixture was vortexed rigorously for 4-5 sec. The tube was placed immediately in a Turner Model TD-20e luminometer and peak light emission was determined and converted to photons using a <sup>63</sup>Ni radioactive light emission calibration standard.

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## RNA Isolation and cDNA Synthesis

Live <u>Renilla reniformis</u> were collected by bottom trawling in shallow waters off Sapelo Island in the state of Georgia at the University of Georgia Marine Institute. The animals were washed thoroughly in fresh seawater, quick frozen in liquid nitrogen, and stored at -80°C. Frozen <u>Renilla</u> were crushed to a fine

powder under liquid nitrogen with a morter and pestle. The powdered tissue was then homogenized with a Waring blender in 4 M guanidine thiocyanate, and total RNA was isolated as described in Chirgwin et al., Biochemistry (1970) 18:5294-5299. Total RNA was then 5 passed over an oligo-dT cellulose column to obtain polyadenylated RNA which was stored as an ethanol precipitate at -20°C. Single and double stranded cDNA were synthesized from poly A+ RNA by modification of the Gubler and Hoffman method, Gubler et al., Gene 10 (1983) 25:263-269, as described below. Following T-4 polymerase blunting and methylation of the cDNAs, synthetic EcoRI linkers were blunt-end ligated. After digestion with EcoRI, the excess linkers were separated from the cDNAs by low-melt agarose gel electrophor-15 esis. Only cDNA's greater than about 650 bp in length were isolated from the low melt gel.

# Construction and Screening of the Agtll Library

Purified cDNA's were ligated into EcoRIdigested \(\lambda\text{gtll.}\) The DNA was then packaged using \(\lambda\)
phage extracts (Gigapack Plus Kit, Strategene).
Several fractions of the packaged library were titered in Y1088 cells; these fractions ranged from 71% to 81%
recombinant phage as determined by the lack of IPTGinducible \(\text{8-galactosidase}\) activity. The total number of recombinant phage was equal to 2.1 \(\text{X 10}^6\) pfu (plaque forming units). The primary library was then amplified in Y1088 cells and stored in 7% DMSO at -80°C. The
titer of the amplified library was 2.5 \(\text{X 10}^7\) pfu/ml and was approximately 65% recombinant.

Two 17-base oligonucleotide probes were synthesized based on amino acid sequence data from isolated peptides derived from V-8 protease digested, native Renilla luciferase. Shown in Figure 4 are the amino acid sequences of the seven V-8 luciferase peptides. The amino acid sequences with the lowest

codon redundancy were selected for synthesis of luciferase oligonucleotide Probe #1 and Probe #2, which are shown highlighted with their derived nucleotide sequences (lower portion of Figure 4). Probe #1 was derived from peptide 7 and contained 32 redundancies, 5 while Probe #2, derived from peptide 1, contained 64 redundancies. The probes were end-labeled with T-4 polynucleotide kinase to high specific activity  $\{4-9\ x$  $10^8$  cpm/ $\mu$ g}. Y1088 cells were infected with enough phage to give 3 X 104 pfu/plate. The infected cells 10 were plated in 6 ml of top agarose onto 150 mm diameter Luria plates containing 50 µg/ml ampicillin. After overnight incubation at 37°C, the plates were chilled at 4°C before performing plaque lifts. To eliminate false positive signals, duplicate nitrocellulose filter 15 plaque replicas were prepared from each master plate. Filters were processed by base treatment followed by neutralization in Tris buffer.

The filters were air dried and baked at 80°C

in vacuo. Prehybridization was for at least 6 hours

37°C in 6X SSC, 50 mM Sodium Phosphate (pH 6.8), 5X

Denhardt's, and 100 ug/ml denatured Herring sperm

DNA. Hybridization was overnight at 37°C in

prehybridization solution with the addition of dextran

sulfate to a final concentration of 10%. The labeled probes were added to the hybridization solution at 1-2

X 10<sup>6</sup> cpm/ml.

Filter washes were done in the presence of tetramethylammonium choride under the conditions

described for a 17-base oligonucleotide in Wood et al.,

Proc. Nat. Acad. Sci. USA (1985) 82:1585-1588. Each duplicate filter was hybridized to both probes in the first round of screening; in subsequent rounds, the duplicate filters were hybridized to either Probe #1 or Probe #2. All cDNA clones were plaque purified after three or four rounds of screening; phage DNA was isolated from each clone on glycerol step gradients as

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described in Grossberger, D., <u>Nuc. Acid. Res.</u> (1987) 15(16):6737.

## DNA Sequence Analysis

5 All DNA sequence analysis was done in the Ml3 vectors mpl8 and mpl9. Single stranded templates were prepared and dideoxynucleotide sequencing was preformed using a Sequenase DNA Sequencing Kit obtained from United States Biochemical Corporation. Sequencing reactions were primed using either the Ml3 universal 10 primer, a primer which hybridized to extraneous Agtll DNA present in some constructs, or the oligonucleotide probes. Sequence data obtained from both ends of the cDNA was analyzed for six base restriction enzyme sites which were used to generate sequencing subclones 15 (Figure 5). In this way, the entire 1.2 kb cDNA was sequenced on both strands (lower portion of Figure All DNA sequences and translated protein sequences were assembled and analyzed using MicroGenie Sequence 20 Software purchased from Beckman.

## Expression in E. coli

The initial luciferase cDNA clone, \RLuc-6, was in the expression vector \( \alpha gtll. \) The clone was 25 ampliflied in Y1088 cells and the high titer stock was used to make lysogens in Y1089. The \alpha RLuc-6 lysogen was then grown in Luria broth plus ampicillin (50 μg/ml) at 37°C. The cells were pelleted, resuspended in TE buffer, and lysed with lysozyme (2 mg/ml). cell debris was then pelleted and the supernatant was 30 assayed for luciferase activity. The 2.2kbp  $\lambda$ RLuc-6 insert which included 1 kb of Agtll lacZ DNA attached to the 3' end was isolated on a low-melt gel and subcloned into the EcoRI/SstI sites of pTZ18R (Pharmacia). This construct, pTZRLuc-1, was used in 35 the expression and purification of recombinant Renilla luciferase.

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## Electrophoretic and Western Analysis

Recombinant luciferase samples were characterized on Commassie-stained SDS-PAGE gels. For Western analysis, the gels were run and transferred to 5 nitrocellulose filters at 30 mA in transfer buffer as described in Burnett, N.W., Analytical Biochemistry (1981)  $\underline{112}$ :195-203. The filters were blocked with 3% BSA and incubated with a 1/1000 dilution of polyclonal rabbit-anti-luciferase antibodies. Next, the filter 10 was washed in TBS and incubated with al/2500 dilution of the secondary antibody, goat-anti-rabbit IgG conjugated to horseradish peroxidase (Bio-Rad). Finally, the filter was washed in TBS and developed with HRP-Color Developing reagent (Bio-Rad). 15

### Emission Spectra

Crude samples of pTZLuc-1 cells were prepared as described previously in this text. The sample was added to 1 ml of luciferase assay buffer; 1 µl luciferin (917 nmol/µl) was added at 1-2 minute intervals to maintain the signal. The bioluminescence emission spectra were obtained and corrected using an on-line computerized SPEX fluorimeter. Multiple spectra were scan averaged to give the final spectrum which was measured from 675 nm to 375 nm.

### Protein Purification

Purification of recombinant Renilla luciferase
from pTZRLuc-1 E. coli extracts was accomplished in
three chromatographic steps. Recombinant luciferase
was purified from pTZRLuc-1 cells as follows: pTZRLucl cells were grown in 20L Luria broth at 37°C at an
OD<sub>600</sub>=0.6 at which time IPTG was added to a final
concentration of 0.5 mM; cells continued to grow
overnight at 30°C. The cells were harvested by
centrifugation, washed in TE, resuspended in 5 ml of 10

mM EDTA (pH 8) per gram of cells, and frozen at -20°C. In a typical purification, 15 to 30 grams of cells were thawed. Lysozyme was added to a final concentration of 4 to 6 mg/ml, and the cells were held on ice for 45 minutes. DNase 1 (10 to 20 mg) was added to the lysate which was sonicated on ice with 1 minute bursts from a Branson Cell Disrupter until 90% of the cells were lysed as evidenced by microscopic examination.

10 The crude material was clarified by centrifugation at 48 X g for 30 minutes and loaded onto the first column. The extract was first run on a DEAE-Cellulose ion-exchange column followed by a G-100 Sephadex gel filtration column and then a Benzoic Acid-Sepharose affinity column. The G-100 column was run in 15 1X Renilla Standard Buffer (1.5 mM Tris, 1.0 mM EDTA pH 7.8). The other columns were run in 1X buffer and were eluted in 10X buffer (DEAE) or Sodium Benzoate in 10X buffer (Benzoic Acid-Sepharose). The first Benzoic Acid column was eluted with 0.1 M sodium benzoate 20 pulse. The second Benzoic Acid column was eluted with a 0 to 0.5 M sodium benzoate gradient. Protein determinations were made by A280 measurements using the extinction coefficient of native luciferase  $\{\epsilon_{280nm}0.1\%$ = 2.1}, by light emission using the specific activity 25 of naive luciferase  $\{1.8 \times 10^{15} \text{ hv sec}^{-1} \text{ mg}^{-1}\}$ , or by Bradford assays as described in Bradford, M., Analytical Biochemistry (1976) 72: 248. Absorption spectra were measured and collected on a Varian Model DMS-100 spectrophotometer. 30

## Isolation and Analysis of ARLuc-6

The primary screen of 1 X 10<sup>6</sup> recombinant phage resulted in the isolation of nine clones which gave identical autoradiographic signals on both replica filters. Of the nine original positives, only five gave signals on the second screening, and only one of

the five hybridized to both probes. The other four hybridized only to Probe #2, which has the greatest sequence redundancy. Restriction enzyme analysis of the five clones revealed that  $\lambda RLuc-3$  and  $\lambda RLuc-8$  were identical and contained a 1.16 kb insert. 5 λRLuc 2, 5, and 6 had insert sizes of 0.8, 2.34 and 1.2 kbp respectively. Only the \alpha RLuc-3 and \alpha RLuc-8 inserts could be exercised from the EcoRI cloning site by EcoRI digestion. The other three inserts had apparently lost one EcoRI linker site; these had to be cut with EcoRI 10 and SstI. Thus, each of these cDNAs contained 1 kb of  $\lambda$ gtll DNA attached at one end. Since only  $\lambda$ RLuc-6 hybridized to both oligonucleotide probes and contained a cDNA of the size necessary to code for an approximately 36 kd protein, it was chosen for DNA 15 sequence analysis.

The 2.2kb EcoRI/SstI fragment, which contained 1 kb of \agtll lac Z DNA, was subcloned into M13 and mpl8 and mpl9 and both strands of the 1.2 kb cDNA were completely sequenced. The entire cDNA sequence is 1196 20 bp, excluding the EcoRI linker (Figure 1). Structurally, it contains a putative initiation codon beginning at nucleotide 10, a stop codon at nucleotide 944, a polyadenylation consensus sequence at nucleotide 1170, and a short polyadenylated tail of seven 25 nucleotides (Figure 1). Also shown underlined in Figure 1 are the two oligonucleotide hybrization sites located at nucleotides 537-554 (Probe #1) and nucleotides 820-836 (Probe #2). The loss of the EcoRI site at the 3' end of the cDNA was confirmed by the 30 sequence analysis.

The cDNA does not contain a stop codon in frame with and upstream from the first initiation codon as an indication that the protein coding region is full length. However, the coding region directs the recombinant systhesis of fully active Renilla luciferase, as discussed below. Translating the cDNA

sequence into an amino acid sequence gave conclusive evidence that the  $\lambda RLuc$ -6 cDNA was a Renilla luciferase cDNA. The translated cDNA sequence contains an open reading frame of 314 amino acids (Figure 2). The first methionine is preceded by three amino acids which may or may not be part of the native protein sequence. If the <u>in vivo</u> translation begins at the first methionine, an open reading frame of 311 amino acids results which codes for a protein of molecular weight ( $M_{\rm r}$ ) -36 Kd.

- The M<sub>r</sub> of native Renilla luciferase has been measured by various methods with values ranging from 33 Kd to 38 Kd. Comparing the amino acid composition of this translated amino acid sequence with that of the previously published native luciferase composition
- shows a very close homology with many identities between the two (Table 2).

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TABLE 2

Amino Acid Composition of Native and Recombinant Renilla reniformis Luciferase

	Amino Acids	Residues (Native)	Residues (Recombinant)
10	Lysine Histidine Arginine Aspartate* Threonine Serine Glutamate** Proline Glycine Alanine Valine Methionine Isoleucine Leucine Tyrosine Phenylalanine Tryptophan	26 10 12 31 9 20 36 17 19 19 23 7 20 23 12	27 10 13 30 6 19 37 18 17 19 23 9 21 22 13
20	Cysteine	7 3	. 8 3

<sup>\*</sup>Aspartate + Asparagine \*\*Gltamate + Glutamine

Native luciferase composition data taken from Matthews et al., Biochemistry (1977) 16: 85-91.

Further evidence that the cDNA does code for luciferase can be seen by comparing the V-8 protease peptide sequences with the translated cDNA sequence (Figure 3). All V-8 peptides were located on the carboxyl-terminal half of the translated coding region beginning at residue 161; several overlapped with one another. Except at one residue, 219, where the cDNA sequence predicts a tryptophan but peptide 6 sequence indicates a leucine at the same position, all peptides matched exactly to regions of the translated sequence. Bunching of the peptides at one end of the

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protein sequence may be due to the amino-terminal half of the native protein being folded in such a way as to be inaccessible to V-8 protease.

# 5 Expression of Recombinant Luciferase in E. coli

The original ARLuc 6 lysogen showed low levels of luciferase activity as determined by light - emission. IPTG induction of ARLuc-6 lysogens led to an approximate 50% decrease in activity. This result was later explained when DNA sequence data revealed that the 3' end of the cDNA was adjacent to the lac Z sequence in Agtll. Therefore, under conditions of IPTG induction, transcription was being forced in the wrong direction with respect to the luciferase cDNA orientation. Presumably, the non-induced luciferase

orientation. Presumably, the non-induced luciferase expression in this construct was due to promoter activity from the left end of Agtll at a site which we have not determined.

The construct pTZRLuc-1 was made to simplify
the isolation of DNA fragments for use as probes in
Southern and Northen analysis (Figure 6). E. coli
cells harboring this plasmid are referred to as
pTZRLuc-1 cells. Similar to \(\chigq\text{tll}\), the pTZ series
"phagemids" contain a polylinker site adjacent to the

lac Z' gene. Expressed genes in this vector could potentially be expressed containing the first 10 to 15 amino acids of β-galactosidase fused to the cDNA translation product. Analysis of pTZRLuc-l cell supernatants for light emission showed that, relative

to λRLuc-6, high levels of luciferase activity were present. Furthermore, induction of pTZRLuc-1 cells with 0.5 mm IPTG led to an increase in luciferase activity of 5-8 fold in crude extracts.

The bioluminescence emission spectrum from

these crude supernatants was identical to the
previously published bioluminescence emission spectrum
for native Renilla luciferase. The wavelength

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distribution of light emission is essentially identical to that reported earlier. The spectrum had an emission maximum ( $\lambda$ max) at 480 nm with a slight shoulder at 400 nm, which presumably corresponded to the luciferase-oxyluciferin complex neutral species excited state.

The pTZRLuc-1 crude supernatants were further characterized by SDS-PAGE. The Coomassie-stained gel contained numerous bands, one of which ran in the vicinity of native luciferase. To confirm that this band was recombinant luciferse, Western analysis was performed using rabbit polyclonal antibodies raised against native Renilla luciferase. The developed Western showed one band that migrated at the same position as native luciferase. No other products indicative of B-galactosidase-luciferase fusion polypeptide were apparent, suggesting that either any putative fusion protein is in too low a concentration to be detected or, more likely, that no fusion protein is made. Though it has not been confirmed by DNA sequence analysis, any pTZRLuc-1 translation products initiating at the  $\beta$ -galactosidase ATG start codon within the first three codons immediately adjacent to the first cDNA start codon may explain why we see IPTG induction of luciferase activity without production of a fusion product.

IPTG induction of recombinant luciferase indicates that its transcription is directed by the lac Z promoter. Since the only candidate ribosome binding site (RBS) is probably positioned too far (18 nucleotides) from the luciferase ATG to be functional, we suspect that a  $\beta$ -galactosidase peptide is being translated to the stop codon immediately adjacent to the luciferase ATG. The translation of a  $\beta$ -galactosidase peptide may facilitate ribosome reintitiation at the luciferase ATG codon (Figure 7). This event could occur if the dinucleotide AG was acting as a RBS for the luciferase cDNA. In this way

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an IPTG inducible, non-fusion luciferase polypeptide could be synthesized. Given the success of recombinant luciferase expression using the pTZ18 vector, which was designed as a multi-purpose in vitro transcription vector rather than an expression vector, it is obvious that other clones can be developed which express luciferase at levels greater than those which we currently obtain.

#### Purification of Recombinant Renilla Luciferase 10

Using the specific activity for native luciferase, we made calculations for the amount of luciferase present in IPTG induced, pTZRLuc-1 crude supernatants and determined that the amount of recombinant luciferase being produced was sufficient to attempt initial purification on a small scale.

In IPTG induced pTZRLuc-l cells, recombinant luciferase represents approximately 12-14% of the total protein in the clarified crude supernatant. Although significant losses of recombinant luciferase were 20 suffered in this initial purification, the amount of starting material and time involved made the loss seem insignificant when compared to the purification of native luciferase. The purification scheme for the recombinant Renilla luciferase is shown in Figure 8; 25 the purification is summarized in Table 3. analysis of the purification steps shows increasing amounts of recombinant luciferase with respect to contaminating protein. The Benzoic Acid-Sepharose luciferase is approximately 99% pure as evidenced by a single band of  $M_r$  equal to 34 Kd. Very slight contamination was noticible on the Coomassie stained gel if more than 20 µg of protein were loaded.

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important feature for the utility of recombinant luciferase in diagnostic applications, many of which require incubation at physiological temperatures.

Purification of recombinant Renilla luciferase has allowed us to determine its amino-terminal sequence. The amino acid sequence of the first 18 residues was determined by Edman degradation. amino acid peak heights of the sequence data indicated that far less protein was actually being sequenced than was initially protein synthesis, it may be that a large percentage of recombinant luciferase is N-formylated at the initiating methionine and thus blocked to the Edman reaction. In spite of this apparent amino-terminal block, enough unblocked species were available that we were able to obtain sequence for the first 18 residues of recombinant luciferase. The amino acid sequence is identical to the translated cDNA sequence from residues 2 through 18 (Figure 7). On cycle 1 of the amino acid sequencing run, we were not able to confirm the presence of a methionine at the first residue as predicted by the cDNA sequence. However, the fact that the two sets of amino acid data are identical from amino acid residue 2 (Threonine) to residue 18 (Proline) strongly supports our assertion that the first predicted methionine in our sequence is acting as the initiation codon in the pTZRLuc-1 construct.

All publications and patent applications mentioned in this specification are herein incorporated by reference to the same extent as if each individual publication or patent application was specifically and individually indicated to be incorporated by reference.

The invention now being fully described, it will be apparent to one of ordinary skill in the art that many changes and modifications can be made thereto without departing from the spirit or scope of the appended claims.

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important feature for the utility of recombinant luciferase in diagnostic applications, many of which require incubation at physiological temperatures.

Purification of recombinant Renilla luciferase has allowed us to determine its amino-terminal 5 sequence. The amino acid sequence of the first 18 residues was determined by Edman degradation. amino acid peak heights of the sequence data indicated that far less protein was actually being sequenced than was initially protein synthesis, it may be that a large 10 percentage of recombinant luciferase is N-formylated at the initiating methionine and thus blocked to the Edman In spite of this apparent amino-terminal block, enough unblocked species were available that we were able to obtain sequence for the first 18 residues 15 of recombinant luciferase. The amino acid sequence is identical to the translated cDNA sequence from residues 2 through 18 (Figure 7). On cycle 1 of the amino acid sequencing run, we were not able to confirm the presence of a methionine at the first residue as 20 predicted by the cDNA sequence. However, the fact that the two sets of amino acid data are identical from amino acid residue 2 (Threonine) to residue 18 (Proline) strongly supports our assertion that the first predicted methionine in our sequence is acting as the initiation codon in the pTZRLuc-1 construct.

All publications and patent applications mentioned in this specification are herein incorporated by reference to the same extent as if each individual publication or patent application was specifically and individually indicated to be incorporated by reference.

The invention now being fully described, it will be apparent to one of ordinary skill in the art 35 that many changes and modifications can be made thereto without departing from the spirit or scope of the appended claims.

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## WHAT IS CLAIMED IS:

- 1. An isolated DNA or RNA molecule, which comprises a nucleotide sequence coding for Renilla luciferase.
  - 2. The molecule of Claim 1, wherein said molecule comprises the luciferase coding sequence:

- 20 GTTGATCAAATCTGAAGAAGGAGAAAAAATGGTTTTGGAGAATAACTTCT
  TCGTGGAAACCATGTTGCCATCAAAAATCATGAGAAAGTTAGAACCAGAA
  GAATTTGCAGCATATCTTGAACCATTCAAAGAGAAAGGTGAAGTTCGTCG
  TCCAACATTATCATGGCCTCGTGAAATCCCGTTAGTAAAAGGTGGTAAAC
  CTGACGTTGTACAAATTGTTAGGAATTATAATGCTTATCTACGTGCAAGT
- GATGATTTACCAAAAATGTTTATTGAATCGGATCCAGGATTCTTTTCCAA
  TGCTATTGTTGAAGGCGCCAAGAAGTTTCCTAATACTGAATTTGTCAAAG
  TAAAAGGTCTTCATTTTTCGCAAGAAGATGCACCTGATGAAATGGGAAAA
  TATATCAAATCGTTCGTTGAGCGAGTTCTCAAAAATGAACAA
- 30 or an equivalent DNA or RNA sequence.
  - The molecule of Claim 2, wherein said molecule is DNA.

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35 4. The molecule of Claim 3, wherein said molecule contains said luciferase sequence.

5. The molecule of Claim 2, wherein said molecule is RNA and contains a sequence equivalent to said luciferase sequence.

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- 6. The molecule of Claim 1, wherein said sequence is preceded by a functional promoter sequence 5' to said sequence.
- 7. The molecule of Claim 6, wherein at least one copy of said sequence is present in a recombinant DNA or RNA vector.
- A genetically engineered microorganism,
   wherein said microorganism comprises the vector of Claim 7.
  - 9. The microorganism of Claim 8, wherein said microorganism is an  $\underline{E}$ .  $\underline{coli}$  strain.

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- 10. An isolated oligonucleotide, comprising at least 10 consecutive nucleotides selected from nucleotide sequence:

TCCAACATTATCATGGCCTCGTGAAATCCCGTTAGTAAAAGGTGGTAAAC

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complementary DNA sequences, and equivalent or complementary RNA sequences.

- 15 ll. The oligonucleotide of Claim 10, wherein said oligonucleotide is labeled with a detectable tag.
- 12. The oligonucleotide of Claim 10, wherein said oligonucleotide comprises at least 14 consecutive20 nucleotides.
- 13. A method of identifying genetic material encoding a coelenterate luciferase gene, comprising; isolating genetic material from a coelenterate organism to form a sample of genetic material, contacting said sample with an oligonucleotide of Claim 10 under hybridizing conditions, and detecting formation of a duplex comprising said oligonucleotide and DNA or RNA present in said sample.
  - 14. The method of Claim 13, where said method comprises a polymerase chain reaction.
- 35 15. A genetically engineered peptide encoded by a nucleotide sequence of Claim 1.

- 16. The peptide of Claim 15, wherein said peptide is non-glycosylated.
- 17. A peptide comprising the amino acid sequence of
  5 Figure 3 or a fragment of said sequence comprising at least
  5 consecutive amino acid residues in which the fragment is
  immunologically reactive with an antibody that specifically
  binds Renilla luciferase, wherein said peptide is free from
  other Renilla peptides.

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FIG. In 30 40 50 AGCTTAAAGA TGACTTCGAA AGTTTATGAT CCAGAACAAA GGAAACGGAT GATAACTGGT 70 80 90 100 110 120 CCGCAGTGGT GGGCCAGATG TAAACAAATG AATGTTCTTG ATTCATTTAT TAATTATTAT 130 140 150 160 170 180 GATTCAGAAA AACATGCAGA AAATGCTGTT ATTTTTTTAC ATGGTAACGC GGCCTCTTCT 210 220 230 TATTTATESC GACATETTET SCCACATATT GAGCCAGTAG CGCGGTGTAT TATACCAGAT 270 CTTATTEGTA TEGECAAATC AGGCAAATCT GETAATGTT CTTATAGGTT ACTTGATCAT 320 330 340 TACAAATATC TTACTGCATG GTTTGAACTT CTTAATTTAC CAAAGAAGAT CATTTTTGTC GGCCATGATT GGGGTGCTTG TTTGGCATTT CATTATAGCT ATGAGCATCA AGATAAGATC 450 460 470 AAAGCAATAG TTCACGCTGA AAGTGTAGTA GATGTGATTG AATCATGGGA TGAATGGCCT 510 520 GATATTGAAG AAGATATTGC GTTGATCAAA TCTGAAGAAG GAGAAAAAAT GGTTTTGGAG 570 580 590 AATAACTICT TCGTGGAAAC CATGTTGCCA TCAAAAATCA TGAGAAAGTT AGAACCAGAA 620 630 640 GAATTTGCAG CATATCTTGA ACCATTCAAA GAGAAAGGTG AAGTTCGTCG TCCAACATTA 690 TCATGGCCTC GTGAAATCCC GTTAGTAAAA GGTGGTAAAC CTGACGTTGT ACAAATTGTT 750 760 AGGAATTATA ATGCTTATCT ACGTGCAAGT GATGATTTAC CAAAAATGTT TATTGAATCG 810 GATCCAGGAT TCTTTTCCAA TGCTATTGTT GAAGGCGCCA AGAAGTTTCC TAATACTGAA 870 880 TTTGTCAAAG TAAAAGGTCT TCATTTTTCG CAAGAAGATG CACCTGATGA AATGGGAAAA 930 TATATCAAAT CGTTCGTTGA GCGAGTTCTC AAAAATGAAC AATAATTACT TTGGTTTTTT 990 1000 1010 ATTTACATIT TICCCGGGIT TAATAATATA AATGTCATTT TCAACAATTT TATTTTAACT 1050 1060 GAATATÍTCA CAGGGAÁCAT TCATATÁTET TGATTAÁÍTT AGCTCGÁÁCT TTACTCTGTC 1110 1120 1130 ATATCATTIT GGAATATTAC CTCTTTCAAT GAAACTTTAT AAACAGTGGT TCAATTAATT 1170 1160 1180 AATATATAT ATAATTACAT TIGTTATGTA ATAAACTCGG TITTATTATA AAAAAA SUBSTITUTE SHEET COLORS

FIG. 2 Ser Leu Lys Met Thr Ser Lys Val Tyr Asp Pro Glu Gln Arg Lys Arg Met Ile Thr Gly Pro Gln Trp Trp Ala Arg Cys Lys Gln Het Asn Val Leu Asp Ser Phe Ile Asn Tyr Tyr Asp Ser Glu Lys His Ala Glu Asn Ala Val Ile Phe Leu His Gly Asn Ala Ala Ser Ser Tyr Leu Trp Arg His Val Val Pro His Ile Glu Pro Val Ala Arg Cys Ile Ile Pro Asp Leu Ile Gly Het Gly Lys Ser Gly Lys Ser Gly Asn Gly Ser Tyr Arg Leu Leu Asp His Tyr Lys Tyr Leu Thr Ala Trp Phe Glu Leu Leu Asn Leu Pro Lys Lys Ile Ile Phe Val Gly His Asp Trp Gly Ala Cys Leu Ala Phe His Tyr Ser Tyr Glu His Gln Asp Lys Ile Lys Ala Ile Val His Ala Glu Ser Val Val Asp Val Ile Glu Ser Trp Asp Glu Trp Pro Asp Île Glu Glu Asp Ile Ala Leu Ile Lys Ser Glu Glu Gly Glu Lys Met Val Leu Glu Asn Asn Phe Phe Val Glu Thr Het Leu Pro Ser Lys Ile Met Arg Lys Leu Glu Pro Glu Glu Phe Ala Ala Tyr Leu Glu Pro Phe Lys Glu Lys Gly Glu Yal Arg Arg Pro Thr Leu Ser Trp Pro Arg Glu Ile Pro Leu Val Lys Gly Gly Lys Pro Asp Val Val Gln Ile Val Arg Asn Tyr Asn Ala Tyr Leu Arg Ala Ser Asp Asp Leu Pro Lys Net Phe Ile Glu Ser Asp Pro Gly Phe Phe Ser Asn Ala Ile Val Glu Gly Ala Lys Lys Phe Pro Asn Thr Glu Phe Val Lys Val Lys Gly Leu His Phe Ser Gln Glu Asp Ala Pro Asp Glu Het Gly Lys Tyr Ile Lys Ser Phe Val Glu Arg Val Leu Lys Asn Glu Gln

20	40	09	3/8	100	120	140	9
		ard TRP	80 GLY	1 1 TYR	12 ASP	14 Tee	160 GLU
			ILE G	LYS T	HIS A	ALA I	ILE G
		TYR L		TYR L			
		SER T	asp leu		VAL GLY	ile, lys	PRO ASP
THR GLY	TXR I			P HIS			
			E PRO	J ASP	PHE	LYS	TRP
TE			ILE	LEU	371	ASP	GLU
X F	ILE		ILE	LEU	ILE	GLN	ASP
LYS ARG	PHE	nsn	CYS	ARG	LYS	HIS	TRP
	SER	HIS GLY ASN	ARG		LYS	GEU	
ARG	ASP	HIS	ALA	SER TYR	PRO	TYR	GLU SER
GLN	LEU	LEU	VAL	GLY	LEU	Ser iyr	ILE (
GLU	VAL	PHE	PRO	ASN	ASN	TYR	VAL ILE
PRO	ASN	ILE	GLU	GLY	LEU	HIS	
ASP	MET	VAL	ILE	SER	LEU	PHE	VAL
VAL TYR	LYS GLN	ALA	HIS	LYS	GLU	ALA	VAL
	LYS	ASN	PRO	GLY	PHE	LEU	SER
LYS	CYS	ALA GLU	VAL VAL	SER	TRP	CXS	ala geu ser val val asp
3A ser	ARG	ALA	VAL	LYS	ALA	ALA	AIA
FIG.3A MET THR SEI	ALA	HIS	HIS	GLY	THR	GLY	HIS
MET	TRP	LXS	ARG	MET	LEU	TRP (	VAL
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MET	GLU PRO	VAL LYS GLY LEU HIS	SEK PHE VAL
DOCID: 4WO   9215673A1>	· · · · · · · · · · · · · · · · · · ·	`SUBSTITUTE	SHEET AND COLUMN ASSESSED.

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- 1) GLU-GLY-ALA LYS-LYS-PHE-PRO-ASN-THR-GLU
- 2) GLU-ARG-UAL-LEU-LYS-ASN-GLU
- 3) GLU-GLY-GLU-LYS-MET-UAL-LEU-GLU
- 4) GLU-ASP-ILE-ALA-LEU-ILE-LYS-SER-GLU
- 5) GLU-SER-ASP-PRO-GLY-PHE-PHE-SER-ASN-ILE-DAL-GLU
- 6) GLU-LYS-GLY-GLU-VAL-ARG-ARG-PRO-THR-LEU-SER-LEU-PRO-ARG-GLU-ILE-PRO-LEU-VAL-LYS-GLY
- 7) GLU-ASN-ASN-PHE-PHE-VAL-GLU

GLU-ASN-ASN-PHE-PHE-UAL

Luciferase Probe #1: GAR-AAT-AAT-TIT-TIT-GT (32 degeneracies) 6 C C C

LYS-LYS-PHE-PRO-ASN-THR

Luciferase Probe #2: RAR-RAR-TIT-CCT-RAT-AC

(64 degeneracies)

6 6 6

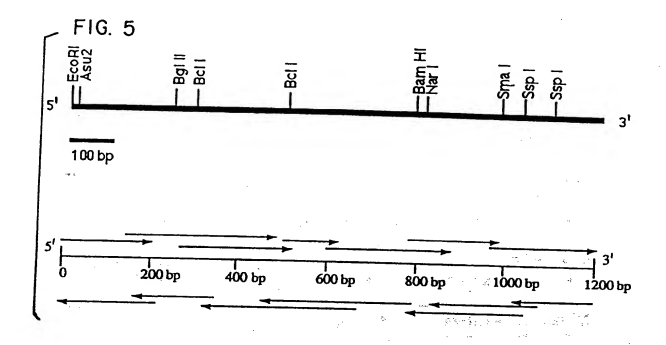
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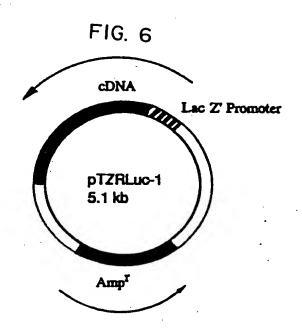
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FIG. 4

SUBSTITUTE SHEET





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pTZRLuc-1

Direction of translation \_\_\_\_\_

lac Z' RBS lac Z' start codon

GTGTGGAATTGTGAGCGGATAACAATTTCACACAGGAAACAGCTATGACCATG

putative RBS \_\_\_\_\_luciferase cDNA "start" codon

ATTACGAATTTAATACGACTCACTATAGGGAATTCAGCTTAAAGATGACTTCG

lac Z' in frame stop codon

AAAGTTTATGATCCAGAA

RBS - ribosome binding site

FIG. 7

SUBSTITUTE SHEET

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## pTZRLuc-1 cells in 10mM EDTA pH8

Freeze thaw once, lysozyme, DNase I, sonication

Centrifugation 48 x g, 30 min

discard pellet

#### DEAE Sephadex

Run in 1X, elute in 10X Standard Buffer, pool activity, concentrate by Amicon Filtration

## G-100 Sephadex

Run in 1X
Standard Buffer,
pool activity,
concentrate by
Amicon filtration

## Benzoic Acid Sepharose 1

Elute with 0.1M Sodium Benzoate in 10X Standard Buffer, dialyze against 1X Standard Buffer

## Benzoic Acid Sepharose 2

Elute with 0 to 0.5M Sodium Benzoate gradient in 10X Standard Buffer, dialyze against 1X Standard Buffer Amicon Filtration

Pure Recombinant Luciferase

FIG. 8

OCID: «WO 9215673A1»

ARTERITATION AND CONTORES

#### INTERNATIONAL SEARCH REPORT

International Antifection No. PCT/US91/01614

I. CLAS	1. CLASSIFICATION OF SUBJECT MATTER 14 several classification symbols, apply, indicate and 5				
Accordin	According to International Patent Classification (IPC) or to both National Classification and IPC				
11' 5' 6	: C12N 9/02, 15/00, 15/11, 15/	53, 15/70. 15/74			
	L.: 435/6, 189, 252.3, 252.33,	320.1; 436/63, 543; 536/	27		
II FIELD	S SEARCHED				
		umentation Searched /			
Classificati		Classification Symnois			
<b></b> .	69.1				
U.	6, 189, 252.3, 25	2.33, 320.1; 436/63, 543;	536/27		
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	Documentation Sourched to	ther than Minimum Documentation			
		nents are Included in the Fields Searched	•		
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APS	Chemical Abstracts/STN, Bios	is/DIALOG databases			
III. DOCI	MENTS CONSIDERED TO BE RELEVANT	·			
Category •	Citation of Document, 11 with indication, where	appropriate, of the relevant passages 12	Relevant to Claim No. 13		
Y	BIOCHEMISTRY, Vol. 16, No. 1	, issued 11 January 1977,	1-9 and		
	MATTHEWS, ET AL., "Purificat	ion and Properties of	15-16		
	Renilla reniformis Luciferas	e", pages 85-91, especi-			
	ally pages 86-89.				
Y	NUCLEIC ACIDS RESEARCH, Vol.		1-9 and		
	25 April 1983, JAYE ET AL.,	"Isolation of a human	15-16		
	anti-haemophiliac factor IX	cDNA clone using unique			
,	52-base synthetic oligonucle				
	the amino acid sequence of be				
	2325-2335, especially pages	2326-2331 and 2334-2335			
	pages .	2320 2331 dild 2334 2333.			
Y	FEBS LETTERS, Vol. 126, No. 1	l issued April 1981	1-9 and		
_	MATSUDA ET AL. "The Primary	Structure of I-1 light			
	MATSUDA ET AL., "The Primary Structure of L-1 Light 15-16 Chain of Chicken Fast Skeletal Muscle Myosin and its				
	Implication", pages 111-113,	soc optime decument			
	implication, pages III-II3,	see entire document.	}		
Y	UO A 97/0220/ (Mapt boy pr	II \ 04 I.m. 1007			
•	WO, A, 87/03304 (McELROY ET A	u.) 04 June 190/,	1-9 and		
	especially pages 16-29.	İ	15-16		
Y	IIC 4 / 05/ 100 /200	1000	1		
r	US, A, 4,956,190 (WOO ET AL.)	23 October 1990,	14		
	especially columns 3 through	17.	]		
		*	1		
		(cont.)			
* Specia	categories of cited documents: 10	T" later document published after to	e international films, date		
"A" document defining the general state of the art weich is not or proprie date and not or Conflict with the implication had					
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"L" document which may throw doubte on propely channels or member an invention step					
Citation or other special reason (as specified) Cannot be Considered to precise an assentive step when the					
"O" document referring to an oral disclosure, use, exhibition or discussional section with the in-more other such that is other means.					
"P" document nutrics of prior to the international file gill the but					
later than the proofst dide claiming 3 cocument member of the Same patient hims.					
IV. CERTIFICATION					
Date of the Actual Completion of the International Search Date of Mahing of this International Search Report					
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	ISA/US	William W. Moore	(vsh)		
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Megor.	Citation of Document, with mulcation, where appropriate, of the resevant passages	Relevant to Claim No.	
à :	THE JOURNAL OF BIOLOGICAL CHEMISTRY, Vol. 254, issued 10 February 1979, WARD ET AL., "An Energy Transfer Protein in Coelenterate Bioluminescence", pages 781-788.	i i	
¥	US, A, 4,983,511 (GEIGER ET AL.) 08 January 1991, especially column 21, line 53, to column 8, line 40.	10-13	
Y	PHOTOCHEMISTRY AND PHOTOBIOLOGY, Vol. 42, No. 5, issued 1985, WEINHAUSEN ET AL. "Luciferases from Different Species of Fireflies are Antigenically Similar", pages 609-611, see entire document.	17	
. <b>A</b> . j. j.	THE JOURNAL OF BIOLOGICAL CHEMISTRY, Vol. 254, No. 3, issued 10 February 1979, CHARBONNEAU ET AL., "CA <sup>2+</sup> -induced Bioluminescence in Renilla reniformis", pages 769-780.	.A. to gra	
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FURTHER INFORM	FURTHER INFORMATION CONTINUED FROM THE SECOND SHEET				
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V. TOBSERVATION	ONE WHERE CERTAIN OF ALMOUSTS SOUND IN				
	ONS WHERE CERTAIN CLAIMS WERE FOUND U				
This international sea	irch report has not been established in respect of certain	claims under Article 17(2) (a) for the	tollowing reasons:		
1. Claim numbers		uired to be searched by this Author	ity, namely:		
	·				
		•			
	•				
2. Claim numbers	haraura they relate to pasts of the interestings.				
	, because they relate to parts of the international a in extent that no meaningful international search can be o	sphication that go not comply with t catried out 13, specifically:	he prescribed require-		
			•		
	. •				
J. Claim numbers	, because they are dependent claims not drafted	in accordance with the second and the	rd sentences of		
PCT Rule 6.4(a).		•			
VI. T OBSERVATIO	ONS WHERE UNITY OF INVENTION IS LACKING	; 2			
			<del></del>		
O3 :	rching Authority found multiple inventions in this internal	tional application as follows:			
Claims 1-14 describe a first Product and Methods of its Making and Use.					
Claims 15-	17 describe a second Product of	that first Method o	f Makina		
and a Metho	od of Use of the latter product.	that IIIst Nethod o	1 Making		
•	and the dated product.	1			
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of the internation	dditional search lens were timely paid by the applicant, this last application. telephone practice	s international search report covers	all searchable claims		
	the required additional search less were timely paid by to				
those claims of H	the international application for which hers were paid, spe	he applicant, this international Sear.	cit tenori covers only		
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J [] No resured additi	linual search less were timely paid by the applicant. Cons	sequently, this international search is	historial progression is a second		
the invention first	mentained in the chams; it is covered by Claim numbers	11			
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4 As all Searchades	chains could be searched watered a field positively are and any additional fee. $\label{eq:conditional}$	Monal fee, the International Society	as a Authority ma not		
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